Fig. 1 SEQ ID NO 1

-496 ACCATAGTCA TCCATATTCA TATTITATAT CATCTCAACT TCTCTTGCCC AAACTTGACC TACTGTTGCA ATTATTITAC ATTTCCTTGG CTCTGTTTTC ATTTATATTT AATTCCAGAA ACCACATCAA GTCTITOCAG AATGAAGTAG AGCATTAAGA AGTAGAGATG TACACACGCA TCTCTAAAAT CAGCCATGCC -296 TACGCAAAGC AGCTTGCACT TAAACACCCA ATACATTTTT CATGATTGTG TTGAAGTGAA GTAACCTAAC CCGTTTTTAT ATCCTTCAAA ATAAGGTGG TAGGAATGCT TTCAGCCCTT TTCAATAGCT TTGATTATCT TGTTTTTGTT ACATCCCTCC TCTTGGTTTG ATCATAGTAG TTACTGTATT TCTTTTTATA 96 ACCTGGTCTG CAAAGGGTAG GCCTTGCAGA CCATTGCAAA GTTGTGACGG CTGTGAGTCA TATTGCTGAA GGTGGAACTC TGAAGGCCAGA CTATCTATGT
AP-1 C/EBP +1 +5 CAAGGCACAA GCTGCTGTTA TATACAACAG AGTGAACTGA GCATCAGTCA GAAAAAGTCT ATGTTTGCAG AAATACAGAT CCAAGACAAA GACAGG ATG +104 GCC ACT GCT GGA AAA GTAAGTGGAACATTT intron 1 (5.6 kb) TTGCCTATTTCTTAG GTT ATT AAA TGC AAA GCA GCT GTG
G T A G K \*\* CTT TOG GAG CAG AAG CAA CCC TTC TCC ATT GAG GAA ATA GAA GTT GCC CCA CCA AAG ACT AAA GAA GTT CGC ATT AAG GTA
L W E Q K Q P F S I E E I E V A P P K T K E V R I K \*\*

20 30 intron 2 (1.0 kb) TGTCTTTGAACACAG ATT TTG GCC ACA GGA ATC TGT CGC ACA GAT GAC CAT GTG ATA

"" I L A T G I C R T D D H V I

40 50 ANA GCA ACA ATG GTG TCC AAG TIT CCA GTG ATT GTG GGA CAT GAG GCA ACT GGG ATT GTA GAG AGC ATT GGA GGA GGG GTG K G T H V S K F P V I V G H E Å T G I V E S I G E G V 60 80 ACT ACA GTG AAA CCA G GTATATGCAGGTGTC intron 3 (0.3 kb) ATTITTTTCCTGTAG GT GAC AAA GTC ATC CCT CTC TTT T V K P \*\* G D K V I P L F CTC CCA CAA TGT AGA GAA IGC AAT GCT TGT CGC AAC CCA GAT GGC AAC CTT TGC ATT AGG AGC GA GTAGGTTTCAGTCAT
L P Q C R E C N A C R N P D G N L C I R S D \*\*

100 110 Intron 4 (0.1 kb) TGATGTATCAAACAG T ATT ACT GGT CGT GGA GTA CTG GCT GAT GGC ACC ACC AGA TTT ACA TGC AAG GGC I T T R F T C K G 117 119 120 ANA CCA GTC CAC CAC TTC ATG AAC ACC AGT ACA TTT ACC GAG TAC ACA GTG GTG GAT GAA TCT TCT GTT GCT AAG ATT GAT K P V H H F H N T S T F T E Y T V V D E S S V A K I D 140 150 160 CAT CCA CCT CCT CCT CAG AAA GTC TGT TTA ATT GCC TGT GGG TTT TCC ACT GGA TAT GGC CCT GCT GTT AAA ACT GGC AAG

D A A P P E K V C L I G C G F S T G Y G A A V K T G X

170

180 CATTCCTCTCCACAG GTC AAA CCT GGT TCC ACT TGC GTC GTC TTT GGC CTG GGA

\*\* V K P G S T C V V F G L G
190 200 GGA GTT GGC CTG TCA GTC ATG AGG GGC TGT AAG TCA GGT GGT GCA TCT AGG ATC ATT GGG ATT GAC CTC AAC AAA GAC AAA G V G L S V I M G C K S A G A S R I I G I D L N K D K 220 TIT GAG AAG GCC ATG GCT GTA GGT GCC ACT GAG TGT ATC AGT CCC AAG GAC TCT ACC AAA CCC ATC AGT GAG GTG CTG TCA
F E K A H A V G A T E C I S P K D S T K P I S E V L S
230 250 ATG ACA GGC AAC AAC GTG GGA TAC ACC TTT GAA GTT ATT GGG CAT CTT GAA ACC ATG GTAAGACCCCAAAAT intron 6
H T G N N V G Y T F E V I G H L E T H \*\*
260 270 CCGTTTTAAACTCAG ATT GAT GCC CTG GCA TCC TGC CAC ATG AAC TAT GGG ACC AGC GTG GTT GTA GGA GTT CCT  $^{\circ}$  I D A L A S C H M N Y G T S V V V G V P 280 290 CCC AAG ATG CTC ACC TAT GAC CCG ATG TTG CTC TTC ACT GGA CGC ACA TGG AAG CGA TGT GTC TTT GGA G GTCAG A K  $\frac{1}{100}$   $\frac{1}{100$ intron 7 (3.2 kb) TGTGCTTATTTGCAG GT TTG AAA AGG AGA GAT GAT GTG CCA AAA CTA GTG ACT GAG TTC  $^{\bullet\bullet}$  G L K S R D D V P K L V T E F 330 CTG GCA AAG AAA TTT GAC CTG GAC CAG TTG ATA ACT CAT GTT TTA CCA TTT AAA AAA ATC AGT GAA GGA TTT GAG CTG CTC L A K K F D L D Q L I T H V L P F K K I S E G F E L L 340 350 360 AAT TCA GGA CAA AG GTAACTGTTTCTTAT INTION 8 (2.3 kb) CATTTTACATTTCAG C ATT CGA ACG GTC CTG ACG TTT TGA N S G Q S \*\* I R T V L T F A 370 374

CATCCAAAGT CCCACGACGT CTGTGTTGTC ATCGTGAACT GCAGTTTCTC TTGTGAGAGT TCCC

Fig. 2

>gi|642473|gb|U16286|HSADHVII1 Human class IV alcohol dehydrogenase (ADH7) gene, exon 1

ACCATAGTCA TCCATATTGA TATTTATAT CATCTCAACT TCTCTTGCCC AAACTTGACC TACTGTGCA ATTATTTAC ATTTCCTTGG
CTCTGTTTTC ATTTATATT AATTCCAGAA ACCACATCAA GTCTTGCAG AATGAAGTAG AGCATTAAGA AGTAGAGATG TACACAGGCA
TCTCTAAAAAT CAGCCATGCCC TAGGCCAAGC ACCTTGCACT TAAACACCCCA ATACATTTT CATGATTATCT TTGAAGTAGA GTAACCTAAC
CCGTTTTATA ATCCTTCAAA ATAAGGTGGA TAGGAATGCT TCAGGCCCTT TTCAATAGCT TTGAATTATCT TGTTTTTGTT AGCTGTATA TCTTTTATATA AGCTGGTCTG CAAAGGGTAG GGCTTGCAGA CCTTGCAGA GTTGCCCCGC
CTGTGGGTCA TATTGCTGAA GGTGGAACTC TGAAGCCAGA CTATCTATGT GAAGGCACAA GCTGCTGTTA TATACAACAG AGTGAACTGA
GCATCAGTCA GAAAAAGTCT ATGTTTGCAG AAATACAGAT CCAAGAGCAAA GACAGGATGG GCACTGCTGG AAAAGTAAGT GGAACATTTC
TGTCCCCTCC TCATCATGAC CTAATGATGT GAGGCCATA CATGAGGGA AACACTGGAT ATGGTGTTTC AGAGCAATTTC
GAGGAATGAA ATGCTTGGTG AGCAGGCATA CAGTGAGGGA AACACTGGAT ATGGTGTTTC AGAGCATTTC
GAGGAATAGTA TATCCAATGT CAGCAAGACA TTGGTGGAAAA ATGCTTGGAGAA AGCCAAATGT TTCCACAAGG ACACTTCAAT TGGCATTATT TGGCATTATT CATGATTTCA TTCCCGGATA
AACACTGGA GCACAGTTGCT CAAAAGTAATTA AACACTTGGA GATATTCCTT
GGCAGGAATGAA TGCTTGGTG AGCCAAATGT TTCCACAAGG ACACTTCAAT TGGCATTATT TGGCATTATT CATAATTTGA TTTCCCGGATA
AG

>qi|642474|qb|U16287|HSADHVII2 Human class IV alcohol dehydrogenase (ADH7) gene, exon 2

TAGGTGACCA TGGATCCATC ATTTAGTCAC TCTGGGCTGC ATGTCTCATG CCTTAAGCAA AGGCACACA AAGATGATT ATAAGGCTTA
TAATACTAGT GTTGATGAA AATCATTCTG GAAAACTTAA AATATTATAT ATGCTCTTC GTTCATATTT ATACTAGTCA GGGAGTCTGT
AAGTTAAAAGG CATAATTCAG ACCATTATA ATACTTGCCT ATTTCTTAGG TTATTAAATG CAAAGCAGCT GTGCTTTGGG ACCAGAAGCA
ACCCTTCTCC ATTGAGGAAA TAGAAGTTGC CCCACCAAAG ACCTAAAAGAAG TCCGATTATA GGTAAGCGTG ACCCTAGAG AACTTAAAGCA
TCCATTACTG GCTCTTATAT ATTGTAAAGG ATCTGGTTGT ACTCGATGTG AAATTAATGA GTTCATTATC C

>gi|642475|gb|U16288|HSADHVII3 Human class IV alcohol dehydrogenase (ADH7) gene, exon 3

AACAATTATA ATTTCTTCAA GAAGTTAAAT ATTCTCCCAA CAGTGAAATG ATCAGTTTGT TGATTGGTGC AATGTTTTTG TCTTTGAACA CAGATTTTGG CCACAGGAAT CTGTCGCACA GATGACCATG TGATAAAAGG AACAATGGTG TCCAAGTTTC CAGTGATTGT GGGACATGAG GCAACTGGGA TTGTAGAGAG CATTGGAGAA GGAGTGACTA CAGTGAAACC AGGTATATGC AGGTGTCAAA CCACAAGTTT GAAATAATTA GGCTTTGATT AGCCTATCAA AGGAAATAGC ACACACTAGG AATTATTAGA GGGATCC

>gi|642476|gb|U16289|HSADHVII4 Human class IV alcohol dehydrogenase (ADH7) gene, exons 4-5

CCTCAACCTT TAGAAGGCAA ACTTACGGTG TTTATAAACC TTAGAATATA TTTTTAAAAG TTTTACCTAT AGTATGGGCT CAATTCACAT TTGTTAATTT CATATTTAA CATTAATGAA CAGCATCTTA TATCATGAT TTTTTCCTGT AGGTGACAAA GTCATCCCTC TCTTTCTGCC ACAATGTAGA GAATGCAATG CTTGTCGCAACCAGTGC AACCTTTGCA TTAGAGAGCGA GTAGGTTTCA GTCATTTTTA CTTTACATTTT TCCTATGCTA ATTTTTGAATTAT TAATAATT AATACGTGTA TTTGATGTAT CAAACACTAT TACTGGTCGT GGAGTACTGG CTGATGGCAC CACCAGATTT ACATGCAAGG CCAAACCAGT CCACCACTTC ATGAACACCA GTACATTTAC CACTGGATA GGCGCTGCTG TTAAAACCTGG CAAGGTAAGA AACCAGGTAGA AACCAGTT TAATTGGCT GTGGGTGATG CAACGAGT CACTGGATA GGCGCTGCTG TAAAAACCTG CAAGGTAAGA AACAGGGTAG CCTAGTTT

>gi|642477|gb|U16290|HSADHVII5 Human class IV alcohol dehydrogenase (ADH7) gene, exon 6

GAGTTCACTC ATGATTTGTG TTTTTAACTG GAGGCCCTTT TCAGGTTTCA CTTTTTGACC CTAACACCTA ACATGTCAA GAACATTCCT CTCCACAGGT CAAACCTGGT TCCACTTGCG TCGTCTTTGG CCTGGGAGGA GTTGGCCTGT CAGTCATCAT GGGCTGTAAG TCAGCTGGTG CATCTAGGAT CATTGGGATT GACCTCAACA AAGACAAATT TGAGAAGGCC ATGGCTGTAG GTGCCACTGA GTGTATCAGT CCCAAGGACT CTACCAAACC CATCAGTGAG GTGCTGTCAG AAATGACCAG CAACAACGTG GGATACACCT TTGAAGTTAT TGGGCATCTT GAAACCATGG TAAGACCCCA AAATTTG

>gi|642478|gb|U16291|HSADHVII6 Human class IV alcohol dehydrogenase (ADH7) gene, exon 7

AACGATCTCC TCCGTTTTAA ACTCAGATTG ATGCCCTGGC ATCCTGCCAC ATGAACTATG GGACCAGCGT GGTTGTAGGA GTTCCTCCAT CAGCCAAGAT GCTCACCTAT GACCCGATGT TGCTCTTCAC TGGACGCACA TGGAAGGGAT GTGTCTTTGG AGGTCAGGAA AGCAAAGCCT CTGGATGGGG AGTGATGGCT TTCACTCTGG TGCTTGGCAA GTGGGAGAAG CCTGTTTCCT CAGGCCTTTC TTCCAAGAAT GAGTATGAAG TGATCT

>qi|642479|qb|U16292|HSADHVII7 Human class IV alcohol dehydrogenase (ADH7) gene, exon 8

AACACAGGGA AAGTCATAGT GCAGGAAAGA GAAGGAATAA ATATAACATA ATAAAAGATA AGGATTATTT AGTAATGTCT AAAGAGAAAA TGTGTGCTTA TTTGCAGGTT TGAAAAGCAG AGATGATGTC CCAAAACTAG TGACTGAGTT CCTGGCAAAG AAATTTGACC TGGACCAGTT GATAACTCAT GTTTTACCAT TTAAAAAAAAT CAGTGAAGGA TTTGAGCTGC TCAATTCAGG ACAAAGGTAA CTGTTTCTTA TGATGA

>gil642480|gb:U16293|HSADHVII8 Human class IV alcohol dehydrogenase (ADH7) gene, exon 9, and complete cds

TTAAACTATC CTTTCTTGAA AGATATGAAA ACAAGTCATT AAAAACTCTC ATTTTACATT TCAGCATTCG AACGGTCCTG ACGTTTTGAG ATCCAAAGTG GCAGGAGGTC TGTGTTGTCA TGGTGAACTG GAGTTTCTCT TGTGAGAGTT CCC

Fig. 3

Fragment number	Size	Contents	Forward and reverse primers
1	450bp	Promoter region, 5' UTR and first exon.	I-F AATCAGCCATGCCTAGGCAAA I-R GGAGGGGACAGAAATGTTCCA
2	405bp	Second exon and parts of flanking introns	2-F GGCTGCATGTCTCATGCCTT 2-R CCAAACATGGGAAGCATCTCC
3	260bp	Third exon and parts of flanking introns	3-F CAGTTTGTTGATTGGTGCAATGTT 3-R TGTGTGCTATTTCCTTTGATAGGCT
4	510bp	Fourth and fifth exon and fourth intron and parts of flanking introns	4-F ATGGGCTCAATTCACATTTGTT 4-R AACTAGCCTACCCTGTTTCTT
5	340bp	Sixth exon and parts of flanking introns	5-F TTCAGGTTTCACTTTTTGACCC 5-R CAAATTTTGGGGTCTTACC
6	260bp	Seventh exon and parts of flanking introns	6-F ATCTCCTCCGTTTTAAACTC 6-R CTCATTCTTGGAAGAAAGGCC
7	265bp	Eighth exon and parts of flanking introns	7-F ACAGGGAAAGTCATAGTGCAGG 7-R TCATCATAAGAAACAGTTAC
8	135bp	Nineth exon and parts of flanking introns	8-F TCCTTTCTTGAAAGATATGAA 8-R CTCCAGTTCACCATGACAACAC

Fig. 4

Name	Frag- ment no.	Loca- tion	Wildtype sequence	Mutated sequence (mutations in bold are underlined)	Restriction enzyme (examples)	Type of mutation
Ml	1	pro- moter	-102 to -88 TTTATAAGTTGGTCT	-102 ω -88 TTTATAAG <u>C</u> TGGTCT	Alu I CviJ I	single nucleotide exchange
M2	1	5'- UTR	+20 to +32 TGTTATATACAAC	+20 to +32 TGTTA <u>C</u> ATACAAC	Mae III	putative TATA-box disrupting
М3	2	second intron	in GDB U16287 nr. 415 to 426 AAATAAGGAGAT	in GDB U16287 nr. 415 to 426 AAATAA <u>GG</u> GGAGAT	-	2bp insertion
M4	3	second intron	in GDB U16288 nr. 81 to 95 TCTTTGAACACAGAT	in GDB U16288 nr. 81 to 95 TCTTTGA <u>G</u> CACAGAT	BsiHKAI Bsp1286I	single nucleotide exchange
M5	3	third exon	in GDB U16288 nr. 205 to 218 GGAGAAGGAGTGAC	in GDB U16288 nr. 205 to 218 GGAGAAG <u>C</u> AGTGAC	(TspR I)	amino acid exchange Gly79Ala
М6	4	fourth intron	in GDB U16289 nr. 274 to 287 ACATTTTTCCTATG	in GDB U16289 nr. 274 to 287 ACATTT <u>G</u> TCCTATG	-	single nucleotide exchange
M7	5	sixth exon	in GDB U16290 nr. 180 to 194 GCATCTAGGATCATT	in GDB U16290 nr. 180 to 194 GCATCTAGAATCATT	Hinf I	silent mutation Arg218Arg

Fig. 5

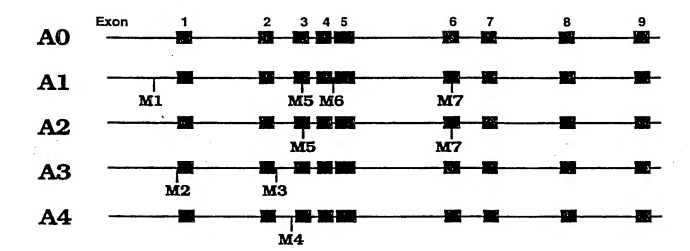


Fig. 6A

	Allele frequency	Number of investigated alleles	Odds ratio	95%CI for the odds ratio	p value
Controls	4.62%	260	-	•	
Non-familial cases	10.23%	88	2.354	0.9565 to 5.796	0.07
Familial cases	28.57%	28	8.267	3.028 to 22.565	0.0002
Patients total	14.66%	116	3.549	1.635 to 7.703	0.0014

Fig. 6B

Allele number	Allele frequency in controls (n=260)	Number of homozygotes in controls (n=130)	Number of homozygotes in PD patients (n=58)	P
· A1	4.62%	0	2	<0.01
АЗ	11.15%	0	3	<0.05